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ARTICLE

Interpenetration Isomers in Isorecticular Amine-tagged Zinc MOFs

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The effect of increasing steric size of pendant amine substituents on structural isorectification has been studied systematically in a series of Zn-MOFs. Linear biphenyl dicarboxylic acids tagged with pendant primary amine (H₂bpdc-NH₂), allylamine (H₂bpdc-NHallyl), diallylamine (H₂bpdc-N(allyl)₂) and dimethylamine (H₂bpdc-NMe₂) groups react with zinc nitrate in DMF to yield a set of interpenetrated MOFs, **WUF-11-14**, respectively, that are structurally akin to IRMOF-9. The allylated amine ligands undergo C-N cleavage reactions under the synthesis conditions, yielding **WUF-12** and **WUF-13** as multivariate MOFs. The single crystal X-ray crystallography on this set of MOFs was not straightforward and we give a salutary account of the difficulties encountered. Gas adsorption measurements combined with surface area calculations provide invaluable support for the crystallographic assignments. The crystallographic analyses reveal subtle differences in the relative positions of the interpenetrating frameworks, and we present a classification system for this type of MOF and analyse related examples available in the literature. CO₂ adsorption measurements revealed that **WUF-14**, which features the strongest Brønsted basic dimethylamine tag group, has the highest capacity, isosteric heat of adsorption, and CO₂/N₂ selectivity.

Introduction

Amino groups are popular choices as functional tags for metal-organic frameworks (MOFs) because the framework topology of the parent MOF can often be replicated using the corresponding aryl amine ligand. This tactic has been used with the most widely investigated aryl amine ligand, 2-aminoterephthalic acid (H₂bdc-NH₂), to produce, *inter alia*, IRMOF-3,^{1, 2} UiO-66-NH₂,³ DMOF-1-NH₂,⁴ UMCM-(3)1-NH₂,⁴ MIL-53-NH₂,^{5, 6} MIL-68(In)-NH₂,⁷ MIL-101(Al)-NH₂,⁸ and MIL-125-NH₂⁹ frameworks by direct synthesis. The underlying motivation for making these materials is often to reap the beneficial effects that the amino functional groups have on the adsorption of CO₂. In addition, the incorporation of an amino group is often part of a strategy for post-synthetic modification (PSM), as they provide nucleophilic sites to subsequently introduce a wide variety of secondary functional groups.¹⁰

The longer congener, 2-amino-[1,1'-biphenyl]-4,4'-dicarboxylic acid (H₂bpdc-NH₂), has received less attention for both direct¹¹⁻¹⁹ and post-synthetic MOF syntheses.^{20, 21} Also, there are only three reports of *N*-alkylated derivatives of H₂bpdc-NH₂.²²⁻²⁴ The purpose of the present work is to detail the structural evolution of zinc(II) MOFs based on the nature of pendant amino substituents on a series of bpdc linkers. The four amino-

functionalised linkers shown in Chart 1 were selected to delineate the impact of steric size and Brønsted basicity. Preparation of an isorecticular MOF series using these linkers would allow the role the functional group plays in the gas adsorption properties to become apparent. There are a limited number of similar studies in the literature.^{2, 25-31} To probe the impacts of the tag groups, we examined CO₂ and N₂ adsorption properties.

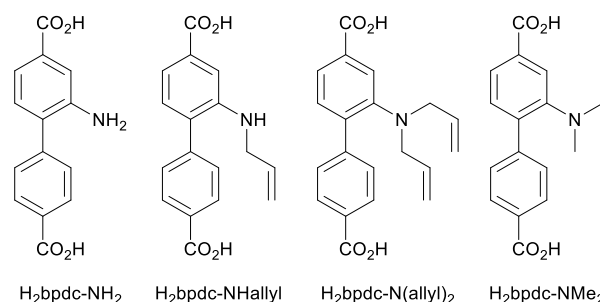


Chart 1 The structures of the amine-tagged biphenyldicarboxylate ligands.

Yaghi and co-workers noted that [1,1'-biphenyl]-4,4'-dicarboxylic acid (H₂bpdc) reacts with zinc(II) nitrate under dilute conditions in *N,N*-dimethylformamide (DMF) solvent to

give crystals with a single, non-interpenetrated $[\text{Zn}_4\text{O}(\text{bpdc})_3]$ framework with a **pcu** topology (IRMOF-10).¹ Curiously, however, follow-up experimental studies on this MOF are rare. Under more concentrated conditions, crystals comprising doubly-interpenetrated **pcu** frameworks are readily formed (IRMOF-9).¹ Although control over interpenetration in a hydroxy-functionalised IRMOF-9/10 system has been demonstrated using special in-situ deprotection conditions,³² a more common approach has been to suppress interpenetration on steric grounds by attaching sufficiently bulky pendant substituents to the biphenyl ligand backbone.^{33–37} Kinetic and coordination modulation methods have also been shown to control interpenetration in other MOFs.^{38,39} It is also possible for bpdc-based zinc IRMOF networks to be partially interpenetrated wherein the second (interpenetrating) lattice is partially occupied.⁴⁰ In this light, the unambiguous assignment of the degree of interpenetration is crucial in the characterisation of IRMOF-9/10 framework analogues. Interpenetration has significant consequences for the positioning and the availability of the functional groups for interactions with guest molecules. Furthermore, the interpenetrating networks may be mutually disposed in various ways, which dictates the shape of the pore environment.

Experimental

Materials and methods

All chemicals used were of analytical grade and purchased from either Sigma Aldrich, VWR Australia or Ajax Finechem Pty Ltd. Synthetic and analytical details for $\text{H}_2\text{bpdc-NH}_2$, $\text{H}_2\text{bpdc-NHallyl}$, $\text{H}_2\text{bpdc-N(allyl)}_2$ and $\text{H}_2\text{bpdc-NMe}_2$ are given in the SI. ^1H NMR and ^{13}C NMR spectra were obtained using a Varian Mercury VX-300-MHz NMR spectrometer operating at 300 MHz for ^1H and 75.5 MHz for ^{13}C , a Bruker Ascend 400 MHz NMR spectrometer operating at 400 MHz for ^1H and 101 MHz for ^{13}C , or a Varian Inova-500-MHz NMR spectrometer, operating at 500 MHz for ^1H and 125 MHz for ^{13}C . ^1H NMR spectra were referenced to the residual *protio* peaks at 2.50 ppm in d_6 -DMSO or 7.27 ppm in CDCl_3 . ^{13}C NMR spectra were referenced to the solvent peaks at 39.6 ppm in d_6 -DMSO or 77.7 ppm in CDCl_3 . For ^1H NMR analysis, MOF samples (~10 mg) were digested by adding 35% DCl in D_2O (2 μL) and d_6 -DMSO (500 μL) and stirring until a solution was obtained.

Simultaneous thermogravimetric and differential thermal analysis (TG-DTA) data were obtained using a Shimadzu DTG-60 instrument fitted with a FC-60A flow rate controller and TA-60WS thermal analyzer. Measuring parameters of 10 $^\circ\text{C}$ per min under nitrogen flow (20 $\text{cm}^3 \text{min}^{-1}$) were used. Powder X-ray diffraction (PXRD) patterns were recorded on a GBC-MMA X-ray diffractometer using $\text{Cu K}(\alpha)$ radiation (1.54180 \AA) with samples mounted on 2.5 cm SiO_2 substrates. Experimental settings in the 2θ angle range of 3–30 $^\circ$ of 0.02 $^\circ$ step size and a scan speed of 1 $^\circ \text{min}^{-1}$ were used. Gas adsorption studies were carried out using a Quantachrome Autosorb MP instrument and high purity nitrogen (99.999%) and carbon dioxide (99.995%)

gases at the Wollongong Isotope Geochronology Laboratory. Surface areas were determined using Brunauer-Emmett-Teller (BET) calculations. Pore size distributions were calculated using the QSDFT kernel for N_2 at 77 K on carbon with slit/cylindrical pores as implemented in the Quantachrome software (v 3.0). The enthalpy of adsorption was calculated by fitting a virial equation to CO_2 isotherms measured at 273 K and 298 K. The Elemental Microanalysis Service at Macquarie University performed microanalyses using a PerkinElmer Elemental Analyzer, Model PE2400 CHNS/O. Each sample was heated at 110 $^\circ\text{C}$ for 2 h and analysed immediately afterward.

Single crystal X-ray diffraction (SCXRD) data were recorded on a Rigaku Spider diffractometer equipped with a MicroMax MM007 rotating anode generator (Cu radiation, 1.54180 \AA), fitted with high flux Osmic multilayer mirror optics, and a curved image-plate detector. Data were collected at 292 K under $d^*\text{TREK}$ and were integrated and scaled and averaged with FS-Process.⁴¹ XPREP was used to determine the space groups and the structures were solved using SHELXS and refined with SHELXL.⁴² Details on the refinement can be found in the ESI to this article. Data are deposited with the Cambridge Structural Database (CCDC 1944583–1944586). Data can be obtained for free from www.ccdc.cam.ac.uk

Synthesis of IRMOF-9- NH_2 [$\text{Zn}_4\text{O}(\text{bpdc-NH}_2)_3$], WUF-11 (WUF, Wollongong University Framework)

$\text{Zn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ (0.346 g, 1.16 mmol) and $\text{H}_2\text{bpdcNH}_2$ (0.100 g, 0.35 mmol) were dissolved in DMF (16 mL) and the solution was placed in an oven for 24 hours at 100 $^\circ\text{C}$. The DMF solution was then exchanged for fresh DMF (2 \times 2 mL) at 100 $^\circ\text{C}$, then at room temperature for CH_2Cl_2 over 3 days, and then for benzene over 3 days. The sample was activated by freeze drying at –53 $^\circ\text{C}$ and 0.09 mbar for 1 hour followed by heating under dynamic vacuum at 120 $^\circ\text{C}$ for 5 hours. Yield 69.8 mg (57%). Anal. Calcd for $\text{C}_{42}\text{H}_{27}\text{N}_3\text{O}_{13}\text{Zn}_4$ [$\text{Zn}_4\text{O}(\text{bpdc-NH}_2)_3$]: C, 48.35; H, 2.61; N, 4.03. Found: C, 48.17; H, 2.36; N, 3.62.

Synthesis and Data for WUF-12-14

Identical synthesis procedures to IRMOF-9- NH_2 were used for all other MOFs.

IRMOF-9-NHallyl (WUF-12): $\text{Zn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ (0.30 g, 1.0 mmol), $\text{H}_2\text{bpdc-NHallyl}$ (0.100 g, 0.34 mmol). Yield 66.5 mg (47 %). Anal. Calcd for $\text{C}_{49.26}\text{H}_{44.68}\text{N}_3\text{O}_{17}\text{Zn}_4$ [$\text{Zn}_4\text{O}(\text{bpdc-NH}_2)_{0.58}(\text{bpdc-NHallyl})_{2.42} \cdot 6\text{H}_2\text{O}$]: C, 47.38; H, 3.93; N, 3.37. Found: C, 47.24; H, 3.41; N, 3.62.

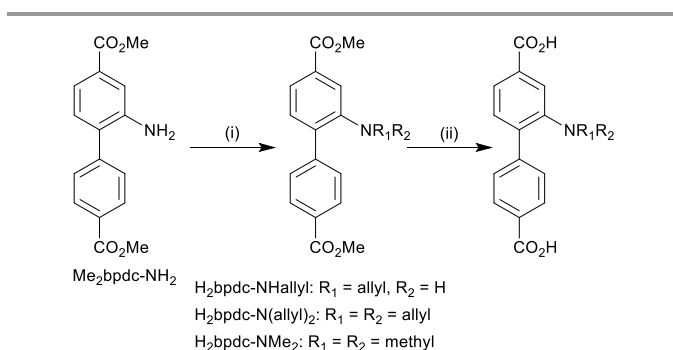
IRMOF-9-N(allyl)₂ (WUF-13): $\text{Zn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ (0.264 g, 0.887 mmol), $\text{H}_2\text{bpdc-N(allyl)}_2$ (0.100 g, 2.96 mmol). Yield 67.1 mg (53%). Anal. Calcd for $\text{C}_{52.62}\text{H}_{49.16}\text{N}_3\text{O}_{17}\text{Zn}_4$ [$\text{Zn}_4\text{O}(\text{bpdc-NHallyl})_{0.54}(\text{bpdc-N(allyl)}_2)_{2.46} \cdot 3.5\text{H}_2\text{O}$]: C, 50.61; H, 3.89; N, 3.37. Found: C, 50.68; H, 3.36; N, 3.84.

IRMOF-9-NMe₂ (WUF-14): $\text{Zn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ (0.313 g, 1.05 mmol), $\text{H}_2\text{bpdc-NMe}_2$ (0.100 g, 0.35 mmol). Yield 69.5 mg (52%). Anal. Calcd for $\text{C}_{48}\text{H}_{45}\text{N}_3\text{O}_{16}\text{Zn}_4$ [$\text{Zn}_4\text{O}(\text{bpdc-NMe}_2)_3 \cdot 3\text{H}_2\text{O}$]: C, 48.80; H, 3.84; N, 3.56. Found: C, 48.99; H, 3.29; N, 4.01.

Results and discussion

Syntheses of the ligands

$\text{H}_2\text{bpdc-NH}_2$ was prepared from $\text{Me}_2\text{bpdc-NO}_2$ by a convenient and high yielding hydrogenation procedure employing ammonium formate and Pd/C, producing $\text{Me}_2\text{bpdc-NH}_2$, followed by hydrolysis in aqueous hydroxide solution (see ESI). Each of the *N*-alkylated ligands were prepared in two-step syntheses starting from $\text{Me}_2\text{bpdc-NH}_2$ via reaction with allyl bromide or methyl iodide, as required, followed by ester cleavage in aqueous hydroxide solution (Scheme 1). Following careful acidification, the ligands were obtained as yellow powders in high yields and characterised by NMR spectroscopy, electrospray ionisation mass spectrometry (ESI-MS) and microanalysis (see ESI).



Scheme 1 Generalised synthetic pathway to the alkylated ligands. Reagents and conditions: (i) allyl bromide/methyl iodide, $\text{K}_2\text{CO}_3/\text{NaHCO}_3$, DMF, 70–90 °C ($\text{Me}_2\text{bpdc-NHallyl}$ 57%; $\text{Me}_2\text{bpdc-N(allyl)}_2$ 85%; $\text{Me}_2\text{bpdc-NMe}_2$ 73%); (ii) NaOH(aq) , THF/MeOH, RT ($\text{H}_2\text{bpdc-NHallyl}$ 97%; $\text{H}_2\text{bpdc-N(allyl)}_2$ 81%, $\text{H}_2\text{bpdc-NMe}_2$ 87%).

Syntheses of the MOFs

MOF syntheses followed a well-established procedure of heating the functionalised bpdc linker with an excess of zinc(II) nitrate in DMF solvent at 100 °C. Using this procedure, IRMOF-9- NH_2 (**WUF-11**), IRMOF-9-NHallyl (**WUF-12**), IRMOF-9-N(allyl) $_2$ (**WUF-13**) and IRMOF-9-NMe $_2$ (**WUF-14**) were obtained as yellow or yellow-brown coloured crystals from $\text{H}_2\text{bpdc-NH}_2$, $\text{H}_2\text{bpdc-NHallyl}$, $\text{H}_2\text{bpdc-N(allyl)}_2$ and $\text{H}_2\text{bpdc-NMe}_2$, respectively. TG-DTA data for the as-synthesised samples (Fig. S9–S12) showed significant endothermic mass losses (47–62%) up to 240 °C, consistent with solvent being driven from the pores of the MOFs. The onset of exothermic decomposition of the frameworks was observed to start around 360 °C.

The as-synthesised MOFs were solvent exchanged with CH_2Cl_2 and dried before digestion in $\text{DCI}/d_6\text{-DMSO}$ for analysis by ^1H NMR spectroscopy (Fig. S1–S4). The spectra show IRMOF-9- NH_2 and IRMOF-9-NMe $_2$ contain solely bpdc- NH_2 and bpdc-NMe $_2$ linkers, respectively; that is, the ligands have been incorporated in the frameworks without modification. However, IRMOF-9-NHallyl and IRMOF-9-N(allyl) $_2$ contain proportions of ligands that have been transformed during MOF synthesis. Specifically, *in situ* C–N bond cleavage occurs at nitrogen centres to expel some of the allyl groups. For example, IRMOF-9-NHallyl also contains bpdc- NH_2 bridging ligands (Fig. S2).

Similarly, IRMOF-9-N(allyl) $_2$ contains bpdc-N(allyl) $_2$ and bpdc-NHallyl bridging ligands. This is clearly seen in the ^1H NMR spectrum of digested IRMOF-9-N(allyl) $_2$ by the presence of signals for the internal alkene of bpdc-N(allyl) $_2$ and bpdc-NHallyl around 5.6 and 5.8 ppm, respectively (Fig. S3). Small amounts of bpdc (~1–3%) were seen in some spectra resulting from cleavage of the $\text{C}_{\text{phenyl}}\text{-N}$ bond rather than one of the $\text{C}_{\text{allyl}}\text{-N}$ bonds. ESI-MS of digested MOF samples supported the C–N cleavage results (Fig. S5–S8). Therefore, IRMOF-9-NHallyl and IRMOF-9-N(allyl) $_2$ are multivariate MOFs.^{43, 44} The ^1H NMR data were used to formulate the framework backbones of the MOFs (Table 1) and show that 18–19% of deallylated ligand is typically incorporated into **WUF-12** and **WUF-13** formed over 24 hours at 100 °C.

Curious as to the origin of the C–N cleavage observed during framework assembly, we carried out control experiments subjecting the respective methyl ester compounds to the conditions of MOF synthesis. No C–N cleavage was detected by ^1H NMR spectroscopy in this set of experiments. Some deallylation of the ester compounds was found, however, when benzoic acid was added to the reaction mixtures. This result suggests reactions of carboxylic acids, perhaps in forming zinc carboxylate clusters, may be responsible, and indicates the dicarboxylic acids play a crucial role in the deallylation phenomenon. ^1H NMR analysis of recovered organic material from the MOF syntheses revealed a similar constitution as that observed in the solids and indicates the deallylation reaction observed here occurs pre-incorporation to the MOF lattice. Syntheses of IRMOF-9-NHallyl and IRMOF-9-N(allyl) $_2$ were then performed at a lower temperature of 80 °C. Crystals started appearing in these reactions after 24 hours and the reactions were stopped at 48 hours. ^1H NMR spectroscopy on digested crystals surprisingly showed the proportion of deallylated ligands rose to 34% and 31%, respectively.

Table 1. Framework formulations determined by ^1H NMR spectroscopy in IRMOF-9-NHallyl and IRMOF-9-N(allyl) $_2$ synthesised at 80 °C and 100 °C.

	Synt hesis Tem p. (°C)	% bpdc NH $_2$	% bpdcN Hallyl	% bpdcN(allyl) $_2$	Framework Formulation
IRM OF- 9- NHa llyl	100	19	81		$[\text{Zn}_4\text{O}(\text{bpdcNH}_2)_{0.58}(\text{bpdcNHallyl})_{2.42}]$
IRM OF- 9- NHa llyl	80	34	66		$[\text{Zn}_4\text{O}(\text{bpdcNH}_2)_{1.2}(\text{bpdcNHallyl})_{1.8}]$
IRM OF- 9- N(all yl) $_2$	100		18	82	$[\text{Zn}_4\text{O}(\text{bpdcNHallyl})_{0.54}(\text{bpdcN(allyl)}_2)_{2.46}]$
IRM OF- 9- N(all yl) $_2$	80		31	69	$[\text{Zn}_4\text{O}(\text{bpdcNHallyl})_{0.93}(\text{bpdcN(allyl)}_2)_{2.07}]$

In situ modification to ligands during framework synthesis are not uncommon, and have even being harnessed for control over ligand incorporation.^{32, 45} Extensive C-N bond cleavage via *N*-demethylation of bdc-NMe $_2$ to bdc-NHMe has previously been observed during the synthesis of IRMOF (80%) and UiO-66 (63%) frameworks,⁴⁶ and the cleavage of a ligand C-C σ -bond has been observed in the synthesis of a zinc MOF.⁴⁷ Notably, we did not see any *N*-demethylation of bpdc-NMe $_2$ in this study (Fig. S4). As mentioned in the introduction, linkers substituted with amino groups represent an important class of MOF linkers, and these results point to an important consideration in the design and use of amino-functionalised ligands.

Structures of the MOFs

The structures of all as-synthesised MOFs were determined by single crystal X-ray diffraction (SCXRD). The analyses confirmed that, as expected, all frameworks are built up from Zn $_4$ O nodes reticulated into cubic **pcu** networks by the linear dicarboxylate linkers, with a general formula of $[\text{Zn}_4\text{O}L_3]$. **WUF-11-14** are doubly-interpenetrated frameworks and are therefore functionalised analogues of IRMOF-9 ($[\text{Zn}_4\text{O}(\text{bpdc})_3]$). Details of the crystallographic refinements can be found in the ESI.

The crystallography, however, was far from straightforward. Crystals of **WUF-11-13** are often highly twinned. The datasets of these crystals give E^2-1 values intermediate between those expected for centrosymmetric (0.968) and noncentrosymmetric (0.736) structures. These datasets could be successfully solved in the space group $P-42_1m$ to give interpenetrated frameworks and refined using a twin model (either as a racemic twin or with a permutative twin matrix $[010/001/100]$). But, the datasets could also be successfully modeled in alternative space groups. For

example, **WUF-11** could be solved in $Pm-3m$, and **WUF-13** in $P-4$. The structures in these alternative space groups are noninterpenetrated i.e. at face value the datasets are compatible with both doubly interpenetrated or noninterpenetrated frameworks! Several observations, however, strongly suggested that the $P-42_1m$ space group, and thus the doubly interpenetrated structures, were correct: (1) the observation of extinction under crossed polarisers precludes the cubic crystal system and thus the putative $Pm-3m$ space group for **WUF-11**; (2) datasets on individual crystals that led to E^2-1 values around 0.9 had systematic absences that pointed to the $P-42_1m$ space group; (3) datasets solved in the alternative space groups led to slightly poorer averaging and refinement statistics (most noticeably R_{int} and R_1) along with significant areas of residual electron density (large Q-peaks) in the framework pores. This electron density could not be successfully assigned to a coherent interpenetrating lattice even with a partial occupancy.

WUF-13 proved to be a particularly difficult case. This MOF was the most likely to be non-interpenetrated as it contains bpdc-N(allyl) $_2$ ligands with the largest steric demand. Crystals of as-synthesised **WUF-13** have the propensity to form as large intergrown crystals that display non-uniform extinction properties, with regions that extinguished polarised light as well as regions that did not, perhaps indicating a delicate balance for interpenetration in this system. A dramatic improvement in optical properties was achieved by solvothermal annealing these crystals at 85 °C in fresh dry DMF for 48 hours. The size of the crystals was followed in this process and typically showed small size reductions of ~2-4% along the two crystal dimensions measured (Fig. S22 and Table S3). This value is very much smaller than is required to convert from a non-interpenetrated to fully interpenetrated phase (25%) via an autocatenation process.⁴⁸ SCXRD analysis on crystals post annealing did, however, show much better diffraction and consistently returned the interpenetrated structure.

The salutary lesson here is that had all twin domains been on the nanoscale, then in all likelihood, extinction and hence twinning would not have been detected. The apparent (non-interpenetrated) SCXRD structures would have been at odds with the gas adsorption data (see **Gas adsorption studies**), which indicates a void volume very much smaller than a noninterpenetrated structure. These datasets serve as a cautionary example for the sole use of SCXRD in distinguishing interpenetrated from non-interpenetrated IRMOF-9-type frameworks. While this distinction is unequivocal in many cases, turning a blind eye to extinction properties under crossed polarisers and poor refinement statistics, may easily have led to the misassignment of interpenetration in some circumstances. The risk would be further exacerbated by the use of crystallographic tools that eliminate residual electron density (such as SQUEEZE in PLATON).

Structural differences exist within this set of interpenetrated structures. By virtue of IRMOF-9-NH $_2$, -NHallyl and -N(allyl) $_2$, crystallising in the $P-42_1m$ space group they share the same positioning between the interpenetrated networks (Fig. 1a, b; ESI Movie 1). In this arrangement, the SBU of the second framework

is situated centrally near one of the square faces of the host framework (Fig. 1b). This arrangement results in closest contacts between the interpenetrated networks occurring around the midpoints of the biphenyl linkers (ca. 3.8 Å between closest phenyl ring carbons). This region is where the amino tag groups reside; however, these groups were not located in difference maps, presumably because of four-site disorder. They were placed in representative sites to complete the crystallographic models and held fixed during structure refinement. The tag groups were implanted at full occupancy so the contributions from the minor dicarboxylates in the multivariate MOFs **WUF-12** and **WUF-13** was not accounted for in the refinements. Rectangular ($\sim 15 \times 7$ Å) and square ($\sim 8 \times 8$ Å) pores are observable along the [010] and [001] directions, respectively (Fig. 1a, b), but the largest pore system in this arrangement ($\sim 10 \times 10$ Å) runs parallel to the [110] direction.

The arrangement of the interpenetrated networks in IRMOF-9-NMe₂ (**WUF-14**) is different. This MOF crystallises in the centrosymmetric space group *R*-3*m* with Zn₄O SBUs of interpenetrating networks related by crystallographic inversion. This arrangement results in positioning the SBU of the second network deeper into the pore bounded by the two networks (Fig. 1c, d; SI Movie 2), rather than near a square face. This results in pores of $\sim 12 \times 14$ Å penetrating the crystal lattice along the [211] and [122] directions. Another much smaller pore of ~ 3 Å exists along the [001] direction. The nitrogen atom of the dimethylamine functional group was located in the difference map but the attached methyl groups were not and these were placed in geometrically reasonable positions and refined with fixed bond lengths and angles.

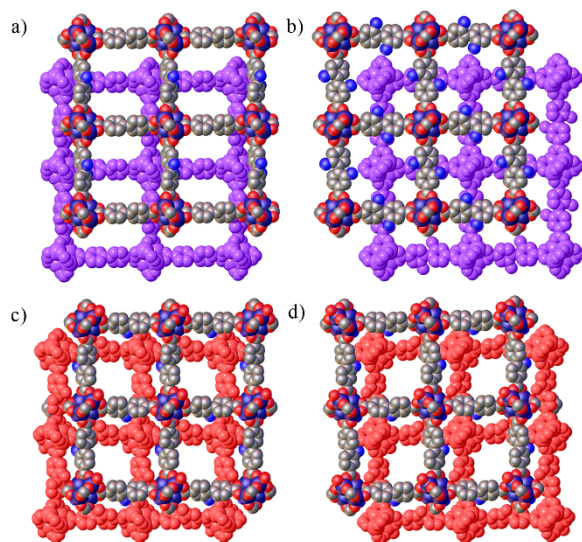


Fig. 1 Orthogonal views of the interpenetration arrangement for IRMOF-9-NH₂ (**WUF-11**), IRMOF-9-NHallyl (**WUF-12**) and IRMOF-9-N(allyl)₂ (**WUF-13**) along a) the b-axis and b) the c-axis; c) and d) orthogonal views for IRMOF-9-NMe₂ (**WUF-14**). Carbon atoms of tag groups and all hydrogen atoms are not shown for clarity

The interpenetration arrangements observed here and in other examples from our labs⁴⁹⁻⁵³ prompted us to classify the modes of interpenetration observed in IRMOF-9 and functionalised

analogues, as determined by SCXRD (Table S2). Three situations for interpenetration are observed (Figure 2). We define Type I as having a SBU located precisely at the centre of the cuboid defined by the host framework and therefore equidistant from all host vertices. Type II has the SBU equidistant from four of the host vertices and can exist from the body centre of the host framework toward the centre of one of the host framework faces. It cannot exist in the face-centred position because of steric clashes with bridging ligands of the host framework. Type III is where the SBU is closest to only one host vertex. Our analysis shows 17 of the 26 interpenetrated IRMOF-9-type structures analysed crystallise as Type III. This arrangement is possible with linker substituents that allow the frameworks to approach closely to maximise noncovalent interactions. There are far fewer frameworks that crystallise in Type II, but this arrangement is observed in three out of the four structures reported here. Although this relative positioning relaxes non covalent interactions between bridging ligands, it provides greater opportunity for larger substituents to point to the central cavity and less chance of encountering steric clashing. This suggests Type II is more suited to larger substituents, such as NHallyl and -N(allyl)₂ found in **WUF-12** and **WUF-13**. Only one system we are aware of crystallises in Type I. This unique system containing bulky phenyl-substituted diazocine substituents allows complementary π - π contacts between the ligands on each framework when the second SBU is located precisely at the centre of the host lattice.

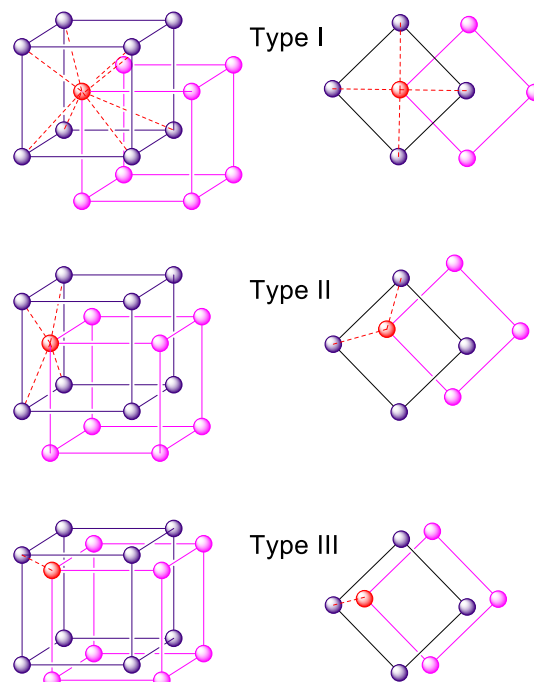


Fig. 2 Perspective and top-down views of the interpenetration Types in IRMOF-9-type structures.

Gas adsorption studies

Figure 3 shows the N₂ isotherms for the MOFs at 77 K, and Table 2 summarises the surface area and pore volume data from

measurements and calculations. Individual adsorption-desorption isotherms are provided in Fig. S28-S31. Each sample shows a Type I isotherm typical of microporous MOF materials. IRMOF-9-NH₂ with the smallest primary amine tag group has the highest apparent BET surface area of 2262 m²g⁻¹, and this compares well with the calculated surface area of 2522 m²g⁻¹. **WUF-12** has an apparent surface area (2095 m²g⁻¹) that is higher than that calculated (1736 m²g⁻¹). This is explained by the calculation model requiring full occupancy NHallyl tags, where the true material has ~19% of these groups cleaved during synthesis to smaller and lighter primary amine groups. **WUF-13**, with the largest diallyl amine functional group, also has a higher apparent surface area (1600 m²g⁻¹) than that calculated (1253 m²g⁻¹) and this is again explained by deallylation. IRMOF-9-NHallyl and IRMOF-9-N(allyl)₂ synthesised at 80 °C have equivalent or higher surface areas (2002 m²g⁻¹ and 1787 m²g⁻¹, respectively) than that synthesised at 100 °C, consistent with that expected based on a greater proportion of deallylated linkers (34% and 31%, respectively). The apparent surface area of IRMOF-9-NMe₂ is 1683 m²g⁻¹ (calc. 2177 m²g⁻¹). The activation of this MOF was particularly difficult. After extensive effort, the best method was supercritical CO₂ drying of the MOF from a benzene solution. Notably, the SCXRD analysis showed this MOF had a water molecule bound to each framework node, and the electron density within the cage-like pore that forms between framework nodes (ESI Movie 3) was modelled as water molecules. We ascribe the difficulties of activating this MOF to successfully removing these water molecules to the presence of and attraction to the -NMe₂ groups that line the cage windows.

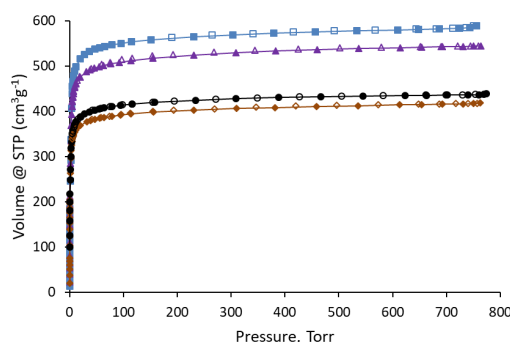


Fig 3. Nitrogen gas adsorption-desorption isotherms at 77 K for IRMOF-9-NH₂ (blue), IRMOF-9-NHallyl (purple), IRMOF-9-N(allyl)₂ (brown) and IRMOF-9-NMe₂ (black). Closed symbols adsorption, open symbols desorption.

The pore size distributions of the MOFs were modelled by DFT calculations. The results show the distributions narrow with increasing size of the amino substituent, consistent with pore constriction by larger functional groups (IRMOF-9-NH₂, 7.5-14 Å; IRMOF-9-NHallyl, 8.5-12 Å; IRMOF-9-N(allyl)₂, 8.5-11 Å) (Fig. S32). These results are in fair agreement with dimensions from SCXRD when considering van der Waals radii. IRMOF-9-NMe₂ shows a distribution tightly clustered between 8 and 11 Å. This is smaller than the dimensions obtained from the SCXRD analyses (~12×14 Å); however, it should be borne in mind that gas adsorption data were recorded at 77 K on activated samples

while the SCXRD data were collected at 292 K on solvated samples. This trend is mirrored in the data for the corresponding pore volumes shown in Table 2, which decrease across the series.

Table 2. Apparent surface areas, pore volumes, isosteric heats of adsorption and CO₂/N₂ selectivity's for the MOFs.

Structure	SA (m ² g ⁻¹) ^a (calc)	Pore Volume (cm ³ g ⁻¹) ^b N ₂ (CO ₂)	$Q_{st}^{CO_2}$ (kJmol ⁻¹) ^c	CO ₂ / N ₂ Selectivity ^d
IRMOF-9-NH ₂	2262 (2522)	0.90 (0.95)	17.5	12.5
IRMOF-9-NHallyl	2095 (1736)	0.84 (0.87)	19	9.5
IRMOF-9-N(allyl) ₂	1600 (1253)	0.64 (0.67)	20	10.0
IRMOF-9-NMe ₂	1683 (2177)	0.67 (0.66)	28	15.1

^a Apparent surface area from BET analysis for N₂ at 77 K. ^b At P/P₀ 0.90 for N₂ at 77 K and at 520 torr for CO₂ at 196 K. ^c From virial fitting calculations. ^d Selectivity from IAST calculations (15% CO₂ and 85% N₂ at 298 K).

We became interested in the effects of desolvation on the Type II MOFs hypothesising that, to achieve greater stabilisation during desolvation, the networks might move relative to the positions determined from the solvated crystal structures. Calculations coupled with SCXRD analyses indicate IRMOF-9 may have considerable plasticity.⁵⁴ We pursued this by measuring the dimensions of crystals of as-synthesised samples as they dried under ambient conditions over 48 hours. IRMOF-9-NH₂ and IRMOF-9-N(allyl)₂ were selected for this part of the study as the MOFs containing the smallest and largest substituents. The measurements are presented in Table S4-S5 and shown in graphical form in Fig. S23-S24 and S26-S27. Crystals of IRMOF-9-NH₂ show size reductions along edge dimensions of ~18-22% over 48 hours with most of the size reduction occurring in the first hour (~10-12%). The size reductions for IRMOF-9-N(allyl)₂ also occur over the first hours but are typically smaller in magnitude (~8-12% over the first hour and ~13-18% over 48 hours). ¹H NMR analysis showed the change in size is congruent with the loss of DMF solvent from the frameworks. After 48 hours, the number of DMF molecules reduced from eight to three and from six to three per formula unit for IRMOF-9-NH₂ and IRMOF-9-N(allyl)₂, respectively. The optical micrographs show crystal morphology is retained during desolvation (see Fig. S25 for an example) yet PXRD studies showed that neither MOF retains sufficient long-range order to give diffraction after more than half an hour in the air. This thwarted further study of the materials by PXRD.

Considering this set of MOFs functionalised with amino tag groups, we were interested in observing their performance in the adsorption of CO₂. The CO₂ sorption data at 196 K (Fig. S33-S38) show that equivalent uptake were obtained compared to the N₂ data at 77 K and corresponding pore volumes were obtained (Table 2). Full adsorption-desorption isotherms recorded from 273 K to 298 K are shown in Fig. S39-S47. The CO₂ adsorption data at 298 K displayed in Fig. 4 is typical of the order observed in the series with IRMOF-9-NMe₂ > IRMOF-9-NHallyl > IRMOF-9-N(allyl)₂ > IRMOF-9-NH₂. The uptake capacities

between 273 K and 298 K are summarised in Table S6 and show IRMOF-9-NMe₂ has the highest uptake at 1 bar with 55 cm³g⁻¹ at 273 K and 31 cm³g⁻¹ at 298 K whereas IRMOF-9-NH₂ displayed the lowest CO₂ uptake with 41 cm³g⁻¹ at 273 K and 23 cm³g⁻¹ at 298 K, despite possessing the highest surface area from N₂ measurements at 77 K. Our measurements agree well with previous GCMC simulations of CO₂ uptake for the IRMOF-9-NH₂ framework.⁵⁵

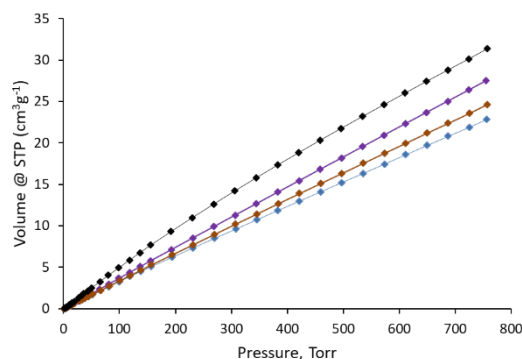


Fig. 4 CO₂ adsorption isotherms recorded at 298 K for IRMOF-9-NH₂ (blue), IRMOF-9-NHallyl (purple), IRMOF-9-N(allyl)₂ (brown) and IRMOF-9-NMe₂ (black).

The selectivity of CO₂/N₂ adsorption was determined by IAST calculations from single-component CO₂ and N₂ isotherms from a theoretical gas mixture consisting of 15 mol% CO₂ and 85 mol% N₂ and is displayed graphically in Fig. S48-S51. The best performing material is again IRMOF-9-NMe₂ with a selectivity of 15.1 and with slightly better performance at 298 K than 273 K. In contrast, IRMOF-9-N(allyl)₂, which also contains a tertiary amine group, shows a relatively poorer performance, with a selectivity of 10.0 at 298 K. IRMOF-9-NHallyl performs similarly to IRMOF-9-N(allyl)₂ although with a slightly lower selectivity of 9.5 while IRMOF-9-NH₂ has a selectivity factor of 12.5 at 298 K.

Recognising the interplay between amine nucleophilicity, basicity and steric profile we sought to probe the strength of the CO₂-framework interaction by calculating Q_{st} values. Fig. 5 shows the results of virial fitting the CO₂ data at 273 K and 298 K for each of the MOFs.

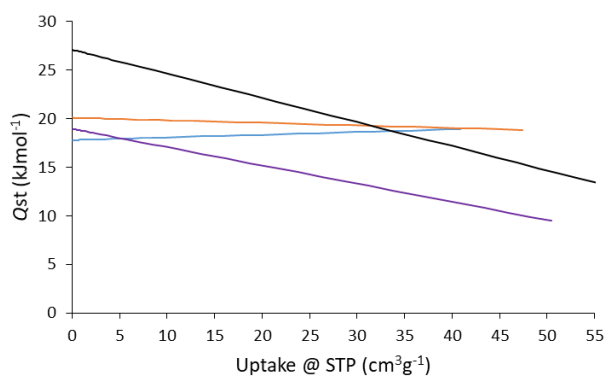


Fig. 5 A plot of the isosteric heats of adsorption of CO₂ for IRMOF-9-NH₂ (blue), IRMOF-9-NHallyl (purple), IRMOF-9-N(allyl)₂ (brown) and IRMOF-9-NMe₂ (black).

The values indicate rather weak physisorption of CO₂ to the frameworks. The tertiary amine MOFs, IRMOF-9-NMe₂ and IRMOF-9-N(allyl)₂, have the highest heats of adsorption. However, the Q_{st} profile for IRMOF-9-NMe₂ reduces comparatively steeply from 27 kJ/mol at low uptake to ~14 kJ/mol at 55 cm³g⁻¹ whereas IRMOF-9-N(allyl)₂ maintains a Q_{st} around 20 kJ/mol out to near 50 cm³g⁻¹. The profile of IRMOF-9-NHallyl also reduces quite steeply while IRMOF-9-NH₂ starts at 17.5 kJ/mol and shows a very small upward trend. In general, the Q_{st} at low coverage follows the same order as the aqueous basicity of the amine: -NMe₂ > -N(allyl)₂ > -NHallyl > -NH₂. As IRMOF-9-NH₂, -NHallyl and -N(allyl)₂ share the same network arrangement, changes in enthalpy of adsorption can be rationalised based on decreasing pore size and increasing amine basicity across these three MOFs. The orientation and availability of the functional groups in the pore environment is also a prime consideration but unfortunately could not be precisely determined by the SCXRD experiments.

In previous work, the enthalpy of adsorption in functionalised IRMOF-9-type compounds was attributed primarily to pore size.⁵⁵ Each of the functionalised IRMOF-9s studied crystallised in Type III with a larger open pore and an inaccessible small pore blocked off by the pendant tag groups, negating effective interaction with CO₂. Although this arrangement is seen here for IRMOF-9-NMe₂ the very strongly basic dimethylamino group leads to a higher enthalpy of adsorption compared to IRMOF-9 despite possessing a larger pore size.

Conclusions

Four interpenetrated amine-containing analogues of IRMOF-9 have been prepared and fully characterised. The dealkylation of secondary and tertiary amines in the solvothermal preparation of the zinc MOFs containing *N*-allyl groups adds to previous observations and suggests this may be a general phenomenon.

The crystal structures of functionalised IRMOF-9 compounds can be classified into three types. Most crystallise with solvent-filled pores as Type III, which offers close inter-framework interactions and one open pore system. The amino-functionalised IRMOF-9-NH₂, IRMOF-9-NHallyl and IRMOF-9-N(allyl)₂ compounds synthesised here crystallise as Type II. This results in close contacts around the inter-ring areas of the biphenyl ligands and leads to multiple types of channels. IRMOF-9-NH₂ and IRMOF-9-N(allyl)₂ show considerable plasticity, as demonstrated by simple desolvation studies. Releasing the full potential of interpenetrated frameworks is likely to come from understanding and harnessing such dynamic behaviour.

In terms of CO₂ adsorption, IRMOF-9-NMe₂ and IRMOF-9-N(allyl)₂ showed the most interesting results. These MOFs contain polar tertiary amines and have smaller pores, indicating electrostatics play a defining role for these systems. The highest CO₂ uptake and selectivity was achieved with the MOF containing the best Brønsted base amongst the functional groups, IRMOF-9-NMe₂.

Conflicts of interest

There are no conflicts of interest to declare.

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Notes and references

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Electronic Supplementary Information (ESI) available: Synthetic procedures, TG–DTA data, descriptions of asymmetric units and additional SCXRD data, ¹H NMR digestion spectra, gas sorption isotherms, virial fitting parameters, IAST plots. See DOI: 10.1039/b000000x/

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